



# **Clinical Edit Criteria Proposal**

Drug/Drug Class Date: Prepared for: Prepared by:	S: Zelnorm® (Tegaserod N August 7, 2002 Missouri Medicaid	/laleate)
New Criteria	<b>a</b>	Revision of Existing Criteria
Executive Su	ummary	
Purpose:	To evaluate and promote pruder indication and duration of use.	nt prescribing of Zelnorm with regards to
Why was this Issue Selected:	irritable bowel syndrome (IBS) in a portion of patients receiving significant drug interactions. IB characterized by symptomatic of structural abnormalities. Commabdominal pain and discomfort function including diarrhea, con	<b>5 5</b>
Program- specific information:	<ul><li>Drug</li><li>Zelnorm 2mg Tabs</li><li>Zelnorm 6mg Tabs</li></ul>	<b>AWP</b> \$172.74/month \$172.74/month
Setting & Population:	NA	
Type of Criteria:	<ul><li>☐ Increased risk of ADE</li><li>☒ Appropriate Indications</li></ul>	<ul><li>☐ Non-Preferred Agent</li><li>☒ Appropriate Utiliztion</li></ul>
Data Sources:	⊠ Only administrative databases	□ Databases + Prescriber- supplied

### **Purpose of Clinical Edit Criteria**

Under the Omnibus Budget Reconciliation Act of 1993, Congress intended Prior Authorization or Prior Approval (PA) programs to control utilization of products that have very narrow indications or high abuse potential. While prescription expenditures are increasing at double-digit rates, payors are also evaluating ways to control these costs by influencing prescriber behavior and guide appropriate medication usage. Clinical Edit criteria, which is different from prior authorization or prior approval programs, assist in the achievement of qualitative and economic goals related to health care resource utilization without placing the entire utilization of a drug in a PA status. Screening the use of certain medications on the basis of clinical appropriateness can reduce costs by requiring evidence of appropriate indications for use, and where appropriate, encourage the use of less expensive agents within a drug class. Clinical Edit criteria can also reduce the risk for adverse events associated with medications by identifying patients at increased risk due to diseases or medical conditions, or those in need of dosing modifications.

### **Setting & Population**

Drug class/Drug for review: Zelnorm<sup>®</sup>

Gender: Female

Diagnosis: IBS, primary bowel symptom constipation

#### **Approval Criteria**

- Patient must be female
- Diagnosis of Irritable Bowel Syndrome
- Primary bowel symptom of constipation
- Therapy approved for no more than 3 annual exacerbations
  - Therapy not to exceed 12 weeks per exacerbation
- Gastroenterologist consult may be required after 2 cycles

#### **Denial Criteria**

- Lack of appropriate diagnostic evidence
- Therapy exceeding approval length

### **Required Documentation**

Laboratory results:	Progress notes:	
MedWatch form:	Other:	



## **Disposition of Edit**

• Denial: Edit 682 "Clinical Edit"

### References

- 1. USPDI, Micromedex, 2003.
- 2. Evidence Based Medicine Analysis: "Zelnorm", UMKC-DIC, January 2004.
- 3. Facts and Comparisons, p. 1165a 1165b, 2004.

